## PATENT SPECIFICATION

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#### (54) PARTICULATE PEROXYGEN COMPOUNDS

(71) We, INTEROX, of 33 Rue Prince Albert, 1050. Brussels, Belgimm, a body corporate, organised and existing under the laws of Belgimm, do hereby declare the invention, for which we pray that a pannt may be grained to us, and the method by which it is to be performed, to be particularly described

in and by the following statement:—
The present invention relates to particulate peroxygen compounds stabilised by coating, to a coating process therefor and to detergent mixtures containing such compounds. is well that percev

known gen compounds can be employed as bleaching compounds in powder detergent mixtures. Sodium perborate tetra-bydrate is usually employed as a bleaching compound in the standard domestic detergents because it is comparatively stable to decom-position in a detargent medium. However, the practice of using cold laundry washing and steeping techniques is becoming increasingly streping techniques is becoming increasingly widespread. Under such conditions, sodium perborate suffers from the disadvantage of disadving too slowly at 20°C. It has been accommended.

It has been suggested that one way of over-coming this disadvantage is to use detergent powders admixed with many other inorganic powers annued with many other morganic peroxygen compounds, particularly the percarbonates, perphosphates and per-cry monosulphates of allali metals, which have suitable dissolution rates. The said peroxygen compounds, particularly the percarbonates, decompose too rapidly, par-ticularly when they are stored in a moist atmosphere. In addition, the other ingredients of the cleaning composition may even trigger decomposition. One method which has been suggested as a way of avoiding this defect

involves coating peroxygen compounds with involves coating perseygen compounts with stabilising substances or protective inorganic films, for example a sodium silicate solution (A. Welter—Br. Pat. 174891 of 26.7.1920) or certain magnesium salts, for example magnesium sulphate heptahydrate. In this case, comparatively large proportions of coating agents must be used in order to ensure adequate

must be used in order to ensure ancquate stability and the granules of stored material show a partial tendency to agglomeration. According to the present invention there is provided, a particulate composition comprising an alkali metal persalt selected from alkali an ansau metal persait senercia from alisali metal percarbonates, perprophosphates, per-tripolyphosphates, persilicates and percor-monosulphates, coated with a mixed salt com-prising sodium carbonate and a compatible sodium salt.

In another aspect of the present invention there is provided a process for at least partistly stabilising an alkali metal persalt elected from alkali metal bonates, perpyrophosphates, pertilicates and percoy-monosulphates to decomposition coverphosphates, persilicates and peroxy-monosculphates to decomposition com-prising coating the persalt with a mixed sait comprising sodium curbonate and a compatible sodium sait. Suimbly the compatible sodium sait is sodium sulphate or sodium curbonate, the mixed sait preferably being sequicarbon-ate. Desirably the mixed sait of sodium saiphate and sodium carbonate has the general formula Na, SO, nNa, CO, wherein n is from

0.3 to 3 and preferably n is from 0.5 to 1.1. In a further aspect of the invention there is provided a process for costing the persalts with the mixed salt comprising depositing the mixed salt comprising depositing the mixed salt come the persalt from a solution containing the constituent parts of the mixed containing the constituent parts of the mixed containing the constituent parts of the mixed persalter conductive to the formation of the mixed salt,

Suitable alkali metal persalts which are normally unstable to water induced a decom-position on storage, and which advantageously may be coated according to the present inven-tion, are preferably the percarbonates, per-pyrophosphates, pertripolyphosphates, and percaymonosulphates.

1.466.799 seen from the phase diagrams of the system.

In the case of the sodium carbonate/sodium

the sodium percarbonate granules, and an 125 aqueous solution to form the appropriate coating agent was sprayed onto the granules through a pneumatic spray located on the wall of the cylinder 11 cms above its base. When

introduction of the solution had been com- 130

Typically sodium percarbonate can be

and 80°C. The nature of the mixed salt will also depend of course on the mole ratio of

the constituents of the mixed salt and can be

coated extremely successfully using a process

2

bicarbonate/water system, the information is available in Kirk Othmer "Encyclopaedia of Chemical Technology" Volume 17 Second coated extremely successfully temp a process
according to the present invention.

Advantageously the coating comprise from
5 0.5% to 20%, by weight based on the presist.
An amount of 0.5%, by weight based on the presist.
An amount of 0.5%, by weight and improved
to emme a tellular participation of the presist.

A tellular participation of the presist of the presis available in Kirk Utimer "Encyclopaedia of Chemical Technology" Volume 17 Second Edition, on page 466. Apparatus in which moving persalt particles appaulus in which moving persur particles may be sprayed, suitably comprises a fluidised bed, a rotary hearth, a rotary drum or other similar recognised devices. It is preferred to 10 weight to ensure a complete coating. A con use a fluidised bed because a more hermetic venient amount is from 3 to 15% by weight and more homogenous coating generally reof coating agent, based on the persalt. and more nomogenous coating generally re-sults, all other conditions of coating being equal, thereby resulting in less coating agent being required to stabilise the persult particles Desirably peroxygen compounds according to the present invention may be used in washing or bleaching compositions. Other standard components may be included in such components. to a given extent, and thus reducing the amount of solvent, usually water, which must sitions, for example bleaching activators, anbe removed by evaporation. Coating using a ionic or non-ionic surfaceants, builders, such as fluidised bed may be carried out continuously carbonates, phosphates, pyrophosphates, tri-polyphosphates, silicates, antiredisposition agents such as carboxymethyl - cellulose, optior barch-wise. If a continuous coating method is used the fluidised bed should preferably be compartmented. Product can be removed from agents such as caromynically a carimose, opin-cal bleaching agents, proteolytic and/or amylolytic enzymes, stabilizers, anti-corrosion agents and the like. Suitable anionic or nonone compartment of such a bed by simply overflowing, or by elutriation via a pipe located at any desired level in the bed. The temperaagents ano the HEC. Sumble amone or non-ionic surfactures include those described by Z. P. Sisley and P. I. Wood in "The Encyclo-peadis of Surface Active Agents" published by Chemical Publishing Ca., New York, Volume 1 in 1961 and in Volume II in 1964, those in French Specification 158987 and those in General Patents Securification 158987 and ture at which simultaneously the persalt is coated, and solvent is evaporated off is of course controlled at below the decomposition temperature of the persait. Naturally the decomposition temperature will vary according to the identity of the persalt, but in general the limit may be set at 100°C. However, in those in German Patent Specification 1594865. In particular, in certain embodiments the case of sodium percarbonate, which is a comparatively heat sensitive product, it is pre-ferred not to exceed approximately 80°C. in particular, in certain embourhers according to the present invention the percey-gen compound satisfies the following test, by losing no more than 15% of active oxygen after four weeks in admirature with a base detergent i.e. a detergent powder which com-Certain embodiments of the present inven-will now be described more fully by way of example. Bassiples 3, 4, 5 and 8 are accord-ing to the present invention. Examples 6 and 7, which describe ceating using respectively 105 neurgent i.e. a accergent powder which contains no peroxygen compound but does contain the other usual ingredients, in a sufficient amount to provide 2% by weight of active oxygen in the mixture. The compositions of and Examples 1 and 2 employing respectively one such base detergent powder is shown in uncoated sodium percarbonate and uncoated Table 2 herein. Storage of the mixture is conducted at 28°C and a relative humidity sodium perborate are present for purposes of 110 comparison only.

In Examples 3 to 8 sodium perbonate was of 70% in cardboard boxes sealed internally and externally by a film of cellulose acetate.

Coating of the persalt particles may be d using a batchwise process in a fluidcoate carried out using any recognised means, and the method employed is not critical but one 50 preferred technique comprises spraying an ised bed. The apparatus used consisted of a cylinder of 15 cm diameter and 77 cm height, fitted at its base with a gas distributor plate pierced aqueous solution of sodium carbonate, with holes of 2 mm diameter, and provided with a pipe bundle for heating using steam maintained at an effective pressure oaf 1 kg/ mining in addition either sodium bicarbona or sodium sulphate onto moving particles of the persalt, and evaporating off water at a 120 55 temperature conducive to formation of a mixed salt, and thereby depositing the mixed First of all the apparatus was charged with 3 kg of homogeneous sodium percarbonate gramles. A stream of air was then passed through the gas distributor plate to fluidise salt on the surface of the persalt. When the mixed salt is to be sodium seswhen the intent sail is to the solution car-quicarbonate or a sodium sulphate/sodium car-60 benate mixed salt the temperature of the solution from which the mixed salt is depo-sited should be maintained at between 30°C

3

### TABLE I

		IABL	45.6			
Coating Conditions	Ex. 3	Ex. 4	Ex. 5	Ex. 6	Ex. 7	Ex. 8
Rate of flow of fluidising gas, m'N./h	38	37	38	31	32	31
Input temperature of fluidising gas, °C	80	102	83	120	120	120
Duration of spraying with solution, minutes	35	40	30	30	30	30
Amount of solution intro- duced, kg	0,47	0,94	0,48	0,60	0,60	0,75
Concentration in the solution in g/kg						
of Na <sub>2</sub> SO <sub>4</sub>	180	180	228		250	_
of Na <sub>2</sub> CO <sub>2</sub>	140	140	85	250	_	172
of NaHCO,	-	_	_	_	_	50
Mole ratio Na <sub>2</sub> CO <sub>3</sub> , Na <sub>2</sub> SO <sub>4</sub>	1,04	1,04	0,5		_	_
Mole ratio Na <sub>2</sub> CO <sub>3</sub> /NaHCO	ł -	-	-		_	2,27
Weight ratio of coating agent to persalt, g/100 g	5	. 10	5	5	5	5
Fluidised bed Temperature	50	50	52	60	60	50
Properties of Coated Percarbonate						
Mean diameter of the granules mm	0,42	0,39	0,42	0,415	0,41	0,41
Free flowing density kg/dm³	1,20	1,15	1,22	1,21	1,19	1,20
Active oxygen content g. kg	134	131	135	135	135	134

Shelf Tests

The following tests were carried out to evaluate the effectiveness of coating percarbonate according to the present invention.

a) Shelf test in small boxes
In this test mixtures were used which connained approximately 2% of active oxygen
comprising 7 g of sodium percarbonate (uncoated or coated) or 10.5 g of sodium perborate (uncoated) and 42 g of a commercial

persalt-free base detergent powder either with or without cazymes having compositions given in Table 2. After homogensision, the mis-tures were placed in cardboard boxes (11.5× 7×2 cm) lined on both sides with a cellulose acctate film resulting in a permeshibity by water of 550 g/m² day. The filled boxes were then stored at 28°C at 70%, relative humidity for 4, 8 and 12 weeks respectively. In a second series of tests further portions of the mixtures were placed in aimlately shade boxes.

which had been coated with a microcrystalline wants into occur control with a microcrystalline wax, to give a permeability by water of  $5 \, g/m^2/dsy$ . These boxes were stored at  $35^{\circ}C$  and  $80^{\circ}g$ , relative himidity for 4 to 8 weeks. After each storage period, the active expensionment of the powder was determined by direct tiration with

and the active oxygen loss relative to the origi-nal active oxygen evaluated. The percentage active oxygen loss of the coated sodium percarbonate was compared with the percentage active oxygen loss of the uncosted sodium percarbonate and of the sodium perborate (stabler product taken as reference) prepared under the same test conditions. The results are expressed in terms of the formula:

in which "T" represented the "active oxygen loss," "PAS" sodium perborate and "PCS" sodium percarbonate. They showed the percentage improvement in the stability emisge improvement in the strenity of sodium percarbonate attributable to coating in relation to the stability of uncoated sodium, perbonate which was taken as a reference be-cause most consumers felt it to be adequate. The active oxygen content was measured by introducing approx. 10 g of washing powder weighed to the nearest 0.01 g in a 750 cc capacity "Riefenmeyer" flasts, adding 100 cc of St H\_SO\_1 100 cc of distilled water and 2 drops of silicone anti-foaming agent. The mixture was then shaken until such time as a homogeneous solution was obtained and then homogeneous solution was obtained and their timed with N/2 KMnO, until a pink colour-ation lasting 30 sec. was obtained. A blank test was also performed on an equivalent amount of washing powder not containing persalt. The active oxygen content of the speci-men, O, expressed in g of oxygen/kg was given by the relationship

$$O=(a-b)\times\frac{1}{2}\times\frac{1000}{2}\times\frac{0.016}{2}$$

#### in which

A. cm<sup>4</sup>, denoted the volume of N/2 KMnO, used for titrating the powder containing persait b, cm<sup>2</sup>, denoted the volume of N/2 KMnO, used for the blank titration

p, g, denoting the weight of sample em-

The results of the foregoing shelf tests are mmarised in the following Tables 3 and 4. summanuscu in the rollowing I ance 3 and 4. These results clearly point to the superiority of stability to decomposition of percarbonate coared with a mixed salt according to the present invention, in comparison with uncontrol percarbonate, or percarbonate coated with only sodium carbonate or sodium sulphate.

TABLE &

IABLE 2.		-
Commercial Powder used in Shelf Tests	Without Enzymes	With Enzymes
Sodium alkylarylsulphonate, g/100 g	24	29
Phesphates - Na, P,O, (*)	. 43	29
Silicates - Na <sub>2</sub> O.2SiO <sub>2</sub> (**) ,,	9	7
Na <sub>2</sub> SO <sub>4</sub> ,,	12	25
н,о "	12	9 .
Enzymes D.U./g		850
(*) total P <sub>2</sub> O <sub>2</sub> expressed in terms of Na <sub>2</sub>	P,O,,	

<sup>(\*\*)</sup> total SiO, expressed in terms of Na, O.2SiO,

In Tables 3 and 4 the persalt is sodium percarbonate unless otherwise stated.

TABLE 3

Results of She	f Tests us	ing Enzym	e-Free De	stergents				
			% Active Oxygen Loss / % improvement in stability					
	Ex.		28°C and 70% RH.		at 35°C	and H.		
Coating of Persalt	No.	4	8	12	4 .	8		
Uncoated	1	13/-	15/-	23/-	17/-	25/-		
Uncoated Sodium Perborate	2	0/-	1/-	2/	4/	6/-		
Na,SO, . 1.04Na,CO,	3	3/77	11/29	17/29	0/131	8/87		
Na_SO 1.04Na_CO_	4	3/17	10/36	16/33	6/85	13/63		
Na SO, . 0.5 Na CO,	5	6/54	12/21	٠.	7/77	13/63		
Na,CO,	6	7/46	13/14 -		10/54	16/47		
Na <sub>2</sub> SO <sub>4</sub>	7	7/46			11/46	17/42		
NaHCO, . Na,CO,	8	5/62						

			% Active % improve	Oxygen L ment in s	oss / . stability	
	_		28°C and		at 35°0	
Coating of Persuit	Ex. No.	4	8	. 12	4 .	8
Uncoated	1	40/-	75/-	88/-	37/-	57/-
Uncoated Sodium Perborate	2	5/-	10/-	32/-	4/-	. 9/-
Na,SO, . 1.04Na,CO,	3_	13/77	41/52	54/61	1/109	12/94
Na,80, . 1.04Na,CO,	4"	14/74	35/62	47/73	7/91	16/85
Na,SO, . 0.5 Na,CO,	. 5	15/71	36/60 <sup>-</sup>	56/57	8/88	16/85
Na <sub>s</sub> CO <sub>s</sub>	6	12/80	44/48	58. 54	14/70 .	30/56
Na,SO,	7	13/77	46/44	. 61./48	14/70	31/54

b) Shelf tests in commercial boxes. In this test mixture were prepared which contained 3.3%, by weight of active oxygen, by mixing 94 grams of tmocated sodium percarbonate having an active oxygen content of 140 g/kg, or 98.5 grams of sodium percarbon-

ate coated with sodium assignicarbonate having an active oxygen content of 134 g/kg with enough persait-free enzyme-free base detergent powder having the composition shown in Table 2 to make up to 400 grams.

After homogenisation the minutes were 15

10

15

placed in carboard boxes of dimensions 20X 14×5.5 cm which had been treated under heat with a micro-crystalline wax. The boxes were then stored at 20° or 35°C and at atmosphere of 80% relative humidity for 2, 4 and 8 weeks, and the tests were carried out in triplicate.

At the end of the selected storage period the active oxygen content of each mixture was measured by tirration of a known mount of mixture dissolved in a known volume of water, against N/10 KMnO. The active oxygen Run in Table 5 following, is expressed as a percentage of the active oxygen present

in the mixture before storage.

In more detail the method of measuring the active oxygen content comprised withthe screw oxygen comment camprised withdrawing a representative sample of 10-20 g,
weighed to the nearest 0.01 g-(P grams)20 and 50 ml of 60°, H<sub>3</sub>SO, dissolved in 1 little
of distilled water 57°C:+2°C in a 2 litre

beaker. The contents were stirred using a one inch SILVERSON LR2 bearing the mark immersed to a depth of 1 cm from the bottom of the beaker, and being rotated at

top speed. "SILVERSON" is a Registered Trade Mark. When the temperature of the solution had fallen to 48-52°C, the weighed sommon has failed to 45-22 C, the wegater sample was added. Stirring continued for 1 minute or longer until all the sample was dis-solved. The solution was allowed to stand for 30 seconds to allow descrution, and 50 ml was transferred to a 250 ml Erlemmeyer flask, and stirred vigorously against standardised N/10 KMnO. The amount of KMnO, solution (T mls) required to reach the end point, a pink colouration lasting at least 15 seconds after addition of 1 more drop of KMnO. was noted. The active oxygen content of the mixture was then expressed using the general formula

30

35

TN 0.8×1050 

wherein N is the actual normality of the KMnO . solution. The results of the tests are summarised in

TABLE 5

	% active oxygen lost during storage at 20°C and at 35°C and							
Coating	80% relative humidity			80% relative humidity				
	2 wks	4 wks	8 Wks	2 wks	4 wks	8 wks		
Uncoated	<del>  -</del> -	5.1	6.4	8.5	9.7	18.1		
5% of sodium sesquicarbonate	-	5.0	5.6	5	5.4	12.3		

These results clearly show the superiority of the products of the invention over the un-50 coated percarbonate.

WHAT WE CLAIM IS:-

1. A particulate composition comprising an alkali metal persalt selected from alkali metal percarbonates, perpyrophosphates, perturboly-phosphates, persilicates and peroxymonosulph-ates, coated with a mixed salt comprising sodium carbonate and a compatible sodium sult

A process for at least partially stabilising an alkali metal persalt selected from alkali metalcarbonates, perpyrophosphates, pertri-polyphosphates, persilicates and peroxymonoates, to decomposition comprising sodium

carbonate and a compatible sodium salt.

3. A compound or process as claimed in claim 1 or 2 wherein the compatible sodium salt is sodium bicarbonate.

4. A compound or process as claimed in

claim 3 wherein the mixed salt is sodium

5. A compound or process as claimed in claim 1 or 2 wherein the compatible sodium

casm 1 or 2 wherein the companies sodium sulphate.

6. A compound or process as claimed in taim 5 wherein the mixed salt has the general formula Na<sub>5</sub>SO<sub>4</sub>.mNa<sub>5</sub>CO<sub>5</sub> wherein n is from 03 m 3.

7. A compound or process as claimed in claim 6 wherein n is from 0.5 to 1.1. claim 6 wherein n is from 0.5 to 1.1.

8. A compound or process as claimed in any preceding claim wherein the persalt is an albali metal percarbonate, perpyrophasphate, pertripolyphosphate or percusymmosulphate.

9. A compound or process as claimed in claim 8 wherein the persalt is sodium percar-

bonate. 10. A compound or process as claimed in

any preceding claim wherein the coating com-prises from 0.5 to 20% by weight of the persalt.

- 11. A compound or process as claimed in claim 10 wherein the casting comprises from 3 to 13% by weight of the personal with 12. A process for conting the peculiar with 12. A process for conting the peculiar with 12. A process for conting the peculiar with 12. A process is claim 12 to 11 comprising deposing the mixed sait onto the persult from a solution containing the constituent parts of the mixed sait in appropriate proportions and at a temperature conductive to the formation of the mixed sait.

  13. A process as claimed in claim 12 where—
  13. A process as claimed in claim 12 where—
  14. The control of the process are formed and it is expected tooms of formed and the process and the control on the process and the process are formed to the process and the process and the process are formed to the process are formed to the process and the process are formed to t
- in the mixed sait is sesquicarbonate formed by evaporation of water from an aqueous solution of sodium carbonate and sodium bicarbonate at a temperature of from 30° to 80°C in a finidised bed.
  - 14. A process as claimed in claim 12 wherein the mixed salt is formed by evaporation of water from an aqueous solution of

- sodium carbonate and sodium sulphate in a mole ratio of from 0.3:1 to 3.0:1 at a tem-perature of from 30 to 80°C in a fluidised bed.
- 15. A process at least partly stabilising a persait substantially as described herein with reference to any one of Examples 3, 4, 5 and

  - A coated persalt whenever produced by a process claimed in any one of claims 2 to 15.
  - 17. A coated persalt substantially as de-scribed herein with reference to any one of
- Examples 3, 4, 5 and 8.

  18. Detergents containing a coated persalt according to any one of claims 1, 3 to 11, 16 and 17.

A. N. CALDWELL, Agent for the Applicants.

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